A Two-Reaction-Zones
Moving-Interface Model for Predicting
$Ca(OH)_2$ Carbonation in Concrete

Michael Böhm
Jörg Kropp
Adrian Muntean

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A Two-Reaction-Zones Moving-Interface Model for Predicting $Ca(OH)_2$-Carbonation in Concrete

Michael Böhm *  Jörg Kropp †  Adrian Muntean ‡

Abstract: Corrosion of steel reinforcement in concrete is essentially controlled by the change of $pH$ near the steel bars. One of the major contributions to a drop in $pH$ is the carbonation of $Ca(OH)_2$ and of other substances near the bars caused by atmospheric $CO_2$ diffusing in the dry parts and reacting in the wet parts of the concrete pores. In this note we propose a prediction model for $Ca(OH)_2$-carbonation. The model is a system of coupled partial differential equations with a moving interface at the carbonation front. It is supplemented by appropriate boundary and interface conditions and allows the prediction of the position of the carbonation front as well as of the concentrations of $CO_2$, rest-$Ca(OH)_2$ and of $CaCO_3$. Special emphasis is laid on the discussion of the reaction near and remote from the carbonation-reaction front. The main tool is the introduction of two separate reaction regions. The formulation of the model allows for 1D to 3D concrete objects. Comparison with penetration curves based on accelerated carbonation experiments as well as with simulated curves show that the model is appropriate for a wide range of concretes.

Key words: Concrete corrosion modeling, carbonation, moving carbonation layer, two-reaction-zones model, coupled partial differential equations, moving-boundary problem.

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1 Introduction

One of the most appreciated features of reinforced concrete structures is their durability, which is endangered by the corrosion of the reinforcing steel. Initially the steel bars are covered by a thin oxide layer which is formed and maintained via its surface by a high alkaline neighbourhood with $pH$ between 12 and 13. The major contributor to the high $pH$ is $Ca(OH)_2$. Depassivation of the steel happens when $Cl^-$ reaches the bars or if the amount of $Ca(OH)_2$ near the bars is reduced by reaction with $CO_2$ - the carbonation, leading to $pH$ well below 9. At a very general level, the $Cl^-$- scenario is pre-dominant under marine

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*ZeTeM, FB3(Mathematik), University of Bremen, Germany, e-mail: mbohm@math.uni-bremen.de
†Institut für Baustofftechnologie, Hochschule Bremen, Germany, e-mail: kropp@fbb.hs-bremen.de
‡ZeTeM, FB3(Mathematik), University of Bremen, Germany, e-mail: muntean@math.uni-bremen.de
conditions, whereas carbonation is dominant under atmospheric exposure and humidity between 0.4 and 0.7. Moreover, there is some interaction between the two mechanisms - with respect to chloride threshold levels and chloride ingress.

This note is a continuation of Ref. [9], in which the concept of a moving carbonation zone and a moving carbonation interface has been introduced and discussed. In the present note we introduce the concept of two reaction zones - one near or at the so-called carbonation front, the second one behind from the first, following the carbonation front.

The aim of this note is to show that this way of modeling captures the typical features of the \( \text{Ca(OH)}_2 \)-carbonation process such as the experimentally verified reaction zone and the fact that after some initial carbonation (in a possibly very narrow reaction zone) the carbonation degree increases gradually until it reaches its maximum.

Experimental results of an accelerated carbonation experiment \((ace)\) will be used for verification of the model.

The main results are summarized in section 5.5.

2 The two-reaction zones model. Introduction

2.1 Generalities

Experimental observation yields the following two typical phenomena during carbonation processes in concrete samples: The concrete sample, \( \Omega \), can roughly be divided in three regions - the uncarbonated part \( \Omega_2 \), a partially carbonated part \( \Omega_{12} \) and an (almost) completely carbonated part \( \Omega_{11} \). The first part is characterized by the complete lack of carbonated substances, the last part contains up to the maximally possible amount of carbonates and in the middle part the concentration of carbonated substances might increase over time. The bulk of the initiating carbonation reaction seems to take place in a very narrow layer \( \Omega_c \subseteq \Omega_{12} \) adjacent to the uncarbonated part \( \Omega_2 \). In most cases this initial reaction does not consume all carbonatable products due to the fact that there may not be a sufficient amount of \( \text{CO}_2 \) to carbonate everything or because some of the carbonatable products are initially encapsulated (which is just a synonym for not yet being available for reaction) and become available for carbonation only after some time. Fig. 1 depicts the geometry of the situation. Fig. 3 depicts possible choices for modeling areas \( \Omega \) in the sample in Fig. 1.

Fig. 5 shows possible scenarios for the advancements of the carbonated zone, the reaction zone and the uncarbonated zone.

Note: In Fig. 5 proportions do not correspond to reality - which is more like

\[
\text{width of } \Omega_c(t) \ll \text{width of } \Omega_1(t) \ll \text{width of } \Omega(t).
\]

In this note we choose the time scale such that \( t = 0 \) corresponds to some time \( \tau_1 > \tau_0 \) in Fig. 5, i.e. in the present situation there is initially a (possibly very small) carbonated part \( \Omega_1(0) \). The reference figure, depicting the initial geometry, is Fig. 6.
Figure 1: Cross section of a circular cylindrical concrete sample $\mathcal{S}$ undergoing carbonation. Atmospheric $CO_2$ and water (humidity) enter $\mathcal{S}$ and react with the carbonatable products in $\mathcal{S}$. A relatively sharp reaction zone $\Omega_\varepsilon$ divides the uncarbonated part $\Omega(t)\backslash\Omega_\varepsilon(t)$ and the rest $\Omega_\varepsilon(t)$ of $\mathcal{S}$. Carbonation continues in $\Omega_\varepsilon(t)$ and, to a lesser extent, in $\Omega_1(t)$. $\Gamma(t)$ is a curve in the center of $\Omega$ which might serve as a substitute for the reaction zone $\Omega_\varepsilon$ (cf. Ref. [9]). All regions change with respect to time $t$. In particular $\Omega_1(t)$ grows and $\Gamma(t)$ moves inwards. The geometrical proportions are exaggerated.

2.2 Specifics

$CO_2$ enters $\Omega$ from the outside through the exterior boundary $\Gamma_{\text{ext}}$, moves in the air-filled parts $\Omega_{\text{ga}}$ of the pores, enters the water-filled parts $\Omega_{\text{gw}}$ of the pores and reacts there with $Ca(OH)_2(aq)$. The reaction is simplified as

$$Ca(OH)_2 + CO_2 \xrightarrow{H_2O} CaCO_3 + H_2O$$

(1)

and is modeled by two (possibly) different reaction rates - a volume rate $\eta_{Rv}$, concentrated on $\Omega_1(t)$, and (a possibly different) volume rate $\eta_{Rv\varepsilon}$ concentrated on $\Omega_\varepsilon(t)$. No $CO_2$ enters the zone $\Omega_2(t)$ which is assumed to be untouched by $CO_2$. Some $Ca(OH)_2(aq)$ is initially present to establish a saturated pore solution. It is consumed by the reaction (1) and can be re-produced by dissolution or other mechanisms. Once $CaCO_3$ is produced by (1) it precipitates instantaneously - in particular it is assumed that there is no $CaCO_3(aq)$-flux. Water is initially present in entire concrete sample $\Omega$. Its concentration is changed due to exterior in- or outflow through $\Gamma_{\text{ext}}$ as well as by the reaction (1). Once $CO_2$ reaches the boundary $\Gamma_r$ (cf. Fig. 2 (b)) of $\Omega$, it ‘tries’ to diffuse into $\Omega_1(t)$. Ultimately one can this consider as the cause of the forward motion of the reaction zone.
2.3 Notations

\( D_{\\text{eff}} \) - effective diffusion coefficient for moisture in \( \Omega_k(t) \), \( k = 1, 2, \epsilon \)

\( D_{\text{ch}} \) - effective diffusion coefficient for \( CO_2(aq) \),

\( D_{\text{ac}} \) - effective diffusion coefficient for \( CO_2(g) \),

\( D_{\text{alk}} \) - effective diffusion coefficient for \( Ca(OH)_2(aq) \),

\( \phi_k \) - volumetric porosities in \( \Omega_k(t) \), \( k = 1, 2 \), with \( \phi_{ks} = 1 - \phi_k \),

\( \phi_w, \phi_a \) - water and air filled, respectively, fraction of the pores in \( \Omega_1(t) \), \( \phi_a + \phi_w = 1 \),

\( \eta_{\text{R}}, \eta_{\text{R,t}} \) - \( \Gamma(t) \)- and \( \Omega_\epsilon(t) \)-concentrated surface- and volume-reaction rates for the reaction (1),

\( P_H \) - mass-transfer coefficient in Henry’s law, \( Q_H \) - exchange factor in Henry’s law,

\( P_{\text{alkdiss}} \) - factor in the dissolution law for \( Ca(OH)_2 \), \( Q_{\text{alkdiss}}, \bar{\eta}_{\text{keq}} \) - equilibrium concentration of \( Ca(OH)_2 \), \( k = 1, 2, \epsilon \) (cf. 3.2.3),

\( m^*_{\text{Ca}(OH)_2(s)}(x) \) - initial (i.e. before dissolution starts) concentration at \( x \) of \( Ca(OH)_2(s) \) which is available for dissolution,

\( S_{\text{alkdiss}} \) - switching factor in the dissolution law,

\( \kappa, \kappa_1 \) - rate constants arising in Arrhenius’ law.

3 The Model. Balance equations, interface, boundary and initial conditions

3.1 Balance equations in \( \Omega_k(t) \), \( k = 1, \epsilon, 2 \)

The geometrical situation is depicted in Fig. 3, Fig. 4 and in Fig. 6. Formally, we attribute to each region \( \Omega_k(t) \), \( k = 1, \epsilon, 2 \), individual source and sink terms (by reaction, dissolution, precipitation and by exchange between wet and dry parts of the pores). These individual
terms will be specified in section 3.2.

- Transport of \( CO_2 \) in the air phase of \( \Omega_1(t) \cup \Omega_\varepsilon(t) \)

\[
\frac{\partial \rho}{\partial t} + \text{div}(\mathbf{j}_\rho) = f_{\text{al}\text{entry}} \quad \text{in} \quad \Omega_1(t) \cup \Omega_\varepsilon(t).
\] (2)

- Transport and reaction of \( CO_2 \) in the wet phase of \( \Omega_1(t) \cup \Omega_\varepsilon(t) \)

\[
\frac{\partial C}{\partial t} + \text{div}(\mathbf{j}_C) = f_{\text{eh}\text{entry}} + f_{\text{rec}} \quad \text{in} \quad \Omega_1(t) \cup \Omega_\varepsilon(t).
\] (3)

- Transport, reaction and dissolution of \( Ca(OH)_2 \) in the wet phase of \( \Omega \)

\[
\frac{\partial \Pi}{\partial t} + \text{div}(\mathbf{j}_{\Pi}) = f_{\text{diss}} + \underbrace{f_{\text{rec}}}_{\text{concentrated on } \Omega_1(t) \cup \Omega_\varepsilon(t)} \quad \text{in} \quad \Omega.
\] (4)

- Transport, reaction, loss by precipitation of \( CaCO_3 \) in the wet phase of \( \Omega_1(t) \cup \Omega_\varepsilon(t) \)

\[
\frac{\partial \phi_{\text{w}} y_m}{\partial t} + \text{div}(\mathbf{j}_{\phi_{\text{w}}}) = f_{\text{reac}} + f_{\text{prec}} \quad \text{on} \quad \Omega_1(t) \cup \Omega_\varepsilon(t).
\] (5)

- The precipitated amount of \( CaCO_3 \) in \( \Omega_1(t) \cup \Omega_\varepsilon(t) \) is obtained from
Figure 4: Another way of choosing the modeling area \( \Omega \) for the cross section of a cylindrical domain. **Geometry of the carbonation process:** \( \Omega := \) reference modeling area section (Compare with Fig. 3). \( \Omega_2 := \) uncarbonated part, \( \Omega_c := \) carbonation-reaction zone adjacent to \( \Omega_1 \), \( \Gamma := \) middle line (or middle surface) of \( \Omega_c, \Gamma_1 \) and \( \Gamma_2 \) - right and left boundary of \( \Omega_c \), \( \Omega_{12} \) and \( \Omega_{12} \) are completely carbonated part and not yet completely carbonated part, \( \Omega_1 := \Omega_{11} \cup \Omega_{12} \). All regions, except \( \Omega \), are time-dependent.

\[
\frac{\partial \phi_{12} \bar{b}_s}{\partial t} + \text{div}(\bar{j}_b) = \phi_{\text{b,prec}} \quad \text{in} \quad \Omega_1(t) \cup \Omega_c(t).
\]

- The total \( CaCO_3 \)-content in \( \Omega_1(t) \cup \Omega_c(t) \) follows from

\[
\frac{\partial \bar{b}}{\partial t} = \phi_{\text{b,prec}} \quad \text{in} \quad \Omega_1(t) \cup \Omega_c(t).
\]

- Transport and generation of \( H_2O \) in \( \Omega \)

\[
\frac{\partial \bar{w}}{\partial t} + \text{div}(\bar{j}_w) = \phi_{\text{w,prec}} \quad \text{in} \quad \Omega.
\]

### 3.2 Specification of reaction, exchange between dry and wet parts of the pores, dissolution and precipitation

#### 3.2.1 Reaction rates

The major purpose of this note is to study the influence of variations of the reaction-rate setting for reaction (1) with respect to the model output (cf. section 5.4). As shown in Ref. [9], the structure (not that much the size!) of the reaction rates influences the output in a significant way. The fact that there is a (possibly very narrow) reaction zone, \( \Omega_c \), does not require two different reaction rates - one on \( \Omega_c \), the other one on \( \Omega_1 \). There is some
information available for the proper choice of a reaction rate - it is based on the analysis of carbonated samples. To our knowledge there is no such information available which refers exclusively to the carbonation-reaction zone $\Omega_\varepsilon$. The common approach is to use one reaction rate for all of $\Omega_1$ and $\Omega_\varepsilon$. 'One' means in particular that all the constants entering the rate formula are the same on $\Omega_1$ and $\Omega_\varepsilon$, respectively. The tacit justification for this is that the reaction mechanism (expressed by the structure of the formula) is the same. Without that assumption one might be tempted to use different rates.

What is the 'right' reaction rate for (1)? Basically, besides doing something completely different, one has three options:

(a) Use reaction rates found by experiment;
(b) Use mass-action law type rates;
(c) Use variations of the mass-action law type rates.

We are not aware of a rich literature with respect to (a). Some authors use (b) - for academic purposes. There are (at least) two more or less common variations as in (c): One
modifies the mass action law by changing the exponents on the concentrations (cf. (12) below), another one (cf. (9)) tries to take the fact into account, that the humidity plays an important role on how fast reaction (1) happens. Using separate rates has its price: One needs more (experimental) a priori information to determine the corresponding constants. The gain is more flexibility, in particular when one deals with very narrow $\Omega_c(t)$ for which we are not aware of experimental data.

In this note we employ the following two reaction rates ((9) and (12));

$$\text{volume reaction rate for (1)} := \eta_{\text{Re}} = \alpha A \exp \left( - \frac{E_0}{RT} \right) f_1 f_2 f_3 f_4 \quad (\text{on } \Omega_1(t)), \quad (9)$$

cf. Refs. [41, 21] also cf. Ref. [46] e.g.), where $\alpha > 0$ is a material constant, $T$ is the absolute temperature, $E_0$ is an activation energy and $A$ is a mean-collision number arising in Arrhenius law, see details in Ref. [2], e.g. $f_1$ is a scenario-switching factor describing the influence of relative humidity $c_w := RH$ on the reaction

$$f_1(c_w) := \begin{cases} 0, & \text{if } 0 \leq c_{w,\text{min}} \\ \frac{5}{2}(c_w - c_{w,\text{min}}), & \text{if } c_{w,\text{min}} < c_w \leq 0.9, \\ 1, & \text{if } 0.9 < c_w \leq 1 \end{cases} \quad (10)$$

where $c_{w,\text{min}} \approx 0.5$. In the accelerated test described in Ref. [45], the relative humidity is fixed to $c_w = 0.65$. The other factors defining $\eta_{\text{Re}}$ are:

$$f_2(\Hat{b}) := 1 - \left( \frac{\Hat{b}}{\Hat{b}_{\text{max}}} \right)^{r_1}, \quad r_1 = \text{const.} > 0, \quad \Hat{b}_{\text{max}} = \text{const.} \quad (\text{Ref. [41]}, \text{one usually considers } r_1 = 1).$$

$$f_3(\Hat{c}) := \frac{\Hat{c}}{\Hat{c}_{\text{max}}} \text{ with } \Hat{c}_{\text{max}} > 0.$$ 

$$f_4(\Hat{h}) := \Hat{h}^{r_2}, \quad r_2 \in [0, 2] \text{ is the newly introduced factor. We remark that if } r_2 = 0, \text{ then (10) is the ansatz used by Wittmann et al (Ref. [21]), Saetta et al (Refs. [41, 42]) and Steffens et al (Refs. [46, 47]), e.g.}$$

This implies the following production rate by reaction in $\Omega_1(t)$:

$$f_{\text{prod}_1} := f_{\text{prod}_2} := -f_{\text{prod}_3} := f_{\text{prod}_4} := \lambda_{\Omega_1(\epsilon)} \eta_{\text{Re}} \quad (11)$$

For the reaction in the layer $\Omega_c(t)$ we propose a simplified ansatz:

$$\epsilon-\text{layer reaction rate for (1)} := \eta_{\epsilon} = \kappa_{\epsilon} A \exp \left( - \frac{E_0}{RT} \right), \quad p, q \geq 0, \quad \kappa_{\epsilon} := \text{rate constant,} \quad (12)$$

where $\kappa_{\epsilon} = \kappa_{\epsilon} \exp \left( - \frac{E_0}{RT} \right)$. The case $p = q = 1$ corresponds to the usual mass-action-law hypothesis. This implies for the production rates by reaction in $\Omega_1(t)$:

$$f_{\text{prod}} := f_{\text{prod}} := -f_{\text{prod}} := f_{\text{prod}} := \lambda_{\Omega_1(\epsilon)} \eta_{\epsilon}. \quad (13)$$

The main reason for introducing $p$ and $q$ is the following: In section 4 we look into the limit case $\epsilon \rightarrow 0$. The mass-action law based rate is formally derived for reactions taking place in volumes of dimensions exceeding the mean free path length of the particles by a factor

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1Within this frame we consider $\Hat{c}_{\text{max}} = \lambda_{\text{CO}_2,\text{out}}$ and $\Hat{b}_{\text{max}} = 0.08324 \text{gcm}^{-3}$. $\Hat{b}_{\text{max}}$ represents the maximum amount of $\text{CO}_2$ which may be consumed by reaction (the degree of hydration is supposed to be about 0.8). $\Hat{c}_{\text{max}}$ is taken to be the concentration of $\text{CO}_2(g)$ from the carbonation chamber.
essentially bigger than 1. Clearly this is not the case if $\varepsilon$ is sufficiently small. Having no theoretical alternative, we try to adapt to the situation by allowing $p, q \neq 1$.

In order to study the influence of the reaction in $\Omega_1(t)$ vs. the one in $\Omega_\varepsilon(t)$, the total production by reaction (1) will be a weighted sum of the individual production rates:

$$ f_{\text{reac}} := \begin{cases} \delta_1 \chi_{\Omega_1(\cdot)} f_{\text{reac}1} + \delta_\varepsilon \chi_{\Omega_\varepsilon(\cdot)} f_{\text{reac} \varepsilon}, & E \equiv \delta, \overline{e}, \overline{w}, \delta_k \geq 0, k = 1, \varepsilon, \end{cases} $$

(14)

where $\alpha$ and $K'_R e$ (cf. (11), (13)) are experimental parameters (cf. Refs. [42, 46, 29]) available only in a certain range. Varying $\delta_k$ ($k = 1, \varepsilon$) can be understood as varying $\delta_1 \alpha$ and $\delta_\varepsilon K'_R e$, respectively. In section 5 we will study the cases $\frac{\delta_\varepsilon}{\delta_1} \ll 1$ and $\frac{\delta_\varepsilon}{\delta_1} \gg 1$.

### 3.2.2 Exchange between dry and wet parts of the pores

In some approaches the reaction (1) is considered as taking place at the interface between the water-filled parts of the pores and the dry part. Then there is no need for an extra pde for $CO_2(aq)$, since the interface approach leads at the pore level (1) to a sink term for $CO_2(g)$ which can directly be coupled with the reaction rate (cf. Refs. [35, 43]). The corresponding result is a modified production-by-reaction rate (at the macroscopic level) incorporating reaction and the idea that the reaction takes place at the interface between $\Omega_{pw}$ and $\Omega_{pu}$. If one thinks of the reaction (1) as taking place in (some part of) $\Omega_{pw}$ then one has to take the exchange of $CO_2(g) \rightleftharpoons CO_2(aq)$ into account. Here this will be done by Henry-law-type exchange terms:

$$ f_{\text{Hentry}} := -f_{\text{Hentry}} = -P_H (Q_H \overline{\varepsilon} - \overline{\varepsilon}) \text{ on } \Omega_1(t) \cup \Omega_\varepsilon(t). $$

(15)

### 3.2.3 Dissolution and precipitation

Dissolution of $Ca(OH)_2$ takes place in all of $\Omega$ as long as there is dissolvable $Ca(OH)_2$. If at time $t = 0$ all dissolvable $Ca(OH)_2$ is in solution, then the corresponding dissolution source $f_{h_diss}, k = 1, 2$ vanishes. Otherwise, the simplest assumption is that of instantaneous (or constant) dissolution, i.e. $f_{h_diss} = Q_{h_diss} \equiv \text{const}$. In this note we model dissolution by the deviation from equilibrium, i.e.

$$ f_{h_diss} = P_{h_diss} (\overline{h}_k - \overline{h}_{eq}) , \; k = 1, 2, $$

(16)

cf. Ref. [28] for a thorough discussion about dissolution in concrete) as long as there is material to dissolve. In order to simplify the notations, we consider that $f_{h_diss} = f_{h_diss}$ in $\Omega_1(t) \cup \Omega_\varepsilon(t)$.

It is reported (cf., e.g., Ref. [11]) that precipitation of $CaCO_3(aq)$ is very fast. We simplify this to instantaneous, i.e.

$$ f_{h_{prec}} := -f_{h_{prec}} := -f_{\text{prec}} \text{ in } \Omega_k, \; k = 1, \varepsilon. $$

(17)

Therefore, the net precipitation rate is

$$ f_{\text{prec}} = \delta_1 \chi_{\Omega_1(\cdot)} f_{\text{reac}1} + \delta_\varepsilon \chi_{\Omega_\varepsilon(\cdot)} f_{\text{reac} \varepsilon} \in \Omega_1(t) \cup \Omega_\varepsilon(t). $$

(18)
3.3 Transport

Transport is modeled by Fick's law, i.e. if \( u_E := [E] \), denotes the concentration of species \( E \), then the flux of \( E \) is

\[
j_E = -D_E \nabla u_E,
\]

where \( D_E \) is the effective diffusion coefficient of species \( E \) in concrete. See, e.g., Refs. [35, 36, 37]. We note that the effective diffusion coefficient \( D_{CO_2} \) might be expressed as \( D_{CO_2} = D_{O_2} f_{diff} \), where \( D_{O_2} \) is the effective diffusion coefficient of \( O_2 \) in concrete and \( f_{diff} \) is a correction factor taking the influence of porosity, temperature and carbonation degree into account (cf. Refs. [3, 46] or Ref. [42] for a simpler setting).

3.4 Two inner-interface conditions at \( \Gamma_r(t) \)

The uncarbonated area is separated from the rest of \( \Omega \) by \( \Gamma_r(t) \). This is the place where \( CO_2 \) might suffer a drop. Furthermore, we assume no \( CO_2 \)-influx into \( \Omega_2(t) \) - since otherwise there could be carbonation going on in \( \Omega_2(t) \). This yields the following Rankine-Hugoniot type boundary condition at \( x + \xi n(x) \in \Gamma_r(t) \) (cf. Ref. [9]):

\[
j_E(x + \frac{\epsilon}{2} n(x), t) \cdot n(x) = E(x + \frac{\epsilon}{2} n(x), t) \hat{s}(x, t) \text{ for all } x \in \Gamma(t), t > 0, E = \overline{\xi, \overline{d}}
\]

The time derivative \( \hat{s}(x, t) \) denotes the speed of progression of the carbonation-reaction layer \( \Omega \epsilon(t) \) in the direction of the normal \( n(x) \) at \( x \in \Gamma(t) \), measured at the middle line (or surface) \( \Gamma(t) \).

We note that, depending on one’s definition of the reaction zone \( \Omega_\epsilon \), the choice

\[
E(x + \frac{\epsilon}{2} n(x), t) = 0, \overline{\xi, \overline{d}}
\]

might be reasonable, as opposed to \( \overline{\xi}(x, t) = 0 \) for \( x \in \Gamma(t) \) in the limit model (\( \epsilon = 0 \)), where the boundary condition (21) is not reasonable, since it implies that there is no reactant available for carbonation at \( \Gamma(t) = \Omega_{\epsilon=0}(t) \). Relation (21) simplifies the flux condition to

\[
j_E(x + \frac{\epsilon}{2} n(x), t) \cdot n(x) = 0 \text{ for all } x \in \Gamma(t), t > 0, E = \overline{\xi, \overline{d}}
\]

A further condition will be needed to complete the model, since the position of \( \Gamma(t) \) is simultaneously to be determined. Balancing the amount produced by carbonation and the amount used for carbonation one obtains (cf. Ref. [9]):

\[
\hat{s}(t) = \frac{1}{\int_{\Omega(t)} n(x + \frac{\epsilon}{2} n(x), t) dx} \epsilon \int_{\Omega(t)} \eta_{HR}(x + \frac{\epsilon}{2} n(x), t) dx \text{ for all } x \in \Gamma(t), t > 0.
\]
3.5 Boundary conditions at the exterior boundary, $\Gamma_{ext}$, and at $x = L$

Boundary conditions on $\Gamma_{ext}$ are needed for all species governed by a pde involving second space derivatives, i.e. for $\vec{c}, \vec{d}, \vec{h}$ and $\vec{w}$, respectively. If there is an oversupply of substance (like for $CO_2(g)$ in accelerated carbonation experiments or for $H_2O$ under heavy rain), or more generally, if one can assume that at a tiny (inner) surface part of the sample the concentrations (or pressures, where applicable) are the same as outside, then one chooses Dirichlet boundary conditions, otherwise Neumann-Robin conditions are the right choice. If $D > 1$, the following setting covers both situations: We divide the boundary $\Gamma_{ext}$ (cf. Fig. 3) in two disjoint parts $\Gamma^{(N)}_{ext}$ and $\Gamma^{(D)}_{ext}$ such that for $E = \vec{c}, \vec{d}, \vec{h}, \vec{w}$:

$$E(x, t) = \lambda_{Eout}(x, t) \text{ for } x \in \Gamma^{(D)}_{ext}, t > 0,$$

$$j_E(x, t) \cdot n(x) = p_E(E(x, t) - \lambda_{Eout}(x, t)) \text{ for } x \in \Gamma^{(N)}_{ext}, t > 0,$$

where $p_E$ denotes a mass-transfer coefficient depending on species $E$ and surface $\Gamma^{(N)}_{ext}$.

In the sequel we restrict ourselves to Dirichlet-boundary conditions at $\Gamma_{ext}$. By symmetry one has to require

$$j_E(x, t) \cdot n(x) = 0, \text{ at } x = L.$$  \hspace{1cm} (26)

3.6 Initial conditions

The initial position of $\Gamma(t)$, denoted by $\Gamma(0)$, and the initial concentrations

$$E(x, 0) = E_0(x), x \in \Omega, \ E = \vec{c}, \vec{d}, \vec{h}, \vec{w},$$

as well as the initial amount of carbonatable $Ca(OH)_2(s)$ (before dissolution starts!), $m^*_n(Ca(OH)_2(s))$, are supposed to be known (cf. appendix).

3.7 The model ($P_{2e}$)

Let $\epsilon > 0$. The balance equations in section 3.1, the production terms in 3.2, the interface, boundary and initial conditions in 3.4 - 3.6 and the flux law in 3.3 will be summarized as model ($P_{2e}$). The solution vector $\overline{X} := (\overline{v}, \vec{c}, \vec{d}, \vec{h}, \vec{w}, \Gamma)$ will be called the model output. If the setting is 1D, then $\Gamma(t)$ can be identified by its position $s(t)$. In this case $\overline{X} := (\overline{v}, \vec{c}, \vec{d}, \vec{h}, \vec{w}, \Gamma, s)$ will be called the model output.
4 The limit case $\epsilon \to 0$ (surface-concentrated reaction + volume reaction)

This short section is a purely academic digression. In Ref. [9] we discussed the case $\epsilon \to 0$, for the following setting (a)-(c):

(a) There is no $Ca(OH)_2$ in $\Omega_1(t)$ and no $CO_2$ in $\Omega_2(t)$,
(b) The carbonation reaction (1) takes place only in $\Omega_\varepsilon(t)$ (as a volume reaction) or
(c) Reaction (1) takes place only on $\Gamma(t)$ (as a surface reaction).

Loosely speaking it turns out that (numerically as well as theoretically) (a) + (c) provides the limit case of (a) + (b) as $\epsilon \to 0$.

In some way there are two limit cases for $\epsilon \to 0$. One models the carbonation reaction as a process taking place solely in the interior of $\Omega_1(t)$, the other one considers (1) as a reaction in $\Omega_1(t)$ plus a reaction concentrated on $\Gamma(t)$.

In order to make this more precise, let $\bar{X}_\varepsilon := (\bar{b}_\varepsilon, \bar{c}_\varepsilon, \bar{d}_\varepsilon, \bar{h}_\varepsilon, \bar{w}_\varepsilon, s_\varepsilon)$ denote the solution of model $(P_{2\epsilon})$ and assume for simplicity $f_{\text{End}} := 0$, keep the structure and constants (with the exception of $\epsilon$) of all material and process functions (porosities, diffusion coefficients, the initial concentrations, the initial position of $\Gamma, \Gamma(0)$, and the boundary setting at $\Gamma_{\text{ext}}$ as well as the the source and sink terms $f_E$, with the exception of the reaction terms for $\Omega_\varepsilon$, $f_{\text{React}}$, fixed.

Let $\Delta V$ be a control volume, $S' := (t, t + \Delta t]$ a time interval. $\int_{S'} \int_{\Delta V \cap \Omega_\varepsilon(\tau)} f_{\text{React}} d\tau d\sigma$ represents the amount of substance associated with the concentration $E$ which is produced by carbonation during $S'$ in $\Delta V$. Under reasonable mathematical assumptions such as uniform boundedness of the concentrations entering $f_{\text{React}}$, it can be shown that there is a limit function $f_{\text{React}}^*$ verifying

$$\int_{S'} \int_{\Delta V \cap \Omega_\varepsilon(\tau)} f_{\text{React}} d\tau d\sigma \xrightarrow{\epsilon \to 0} \int_{S'} \int_{\Delta V \cap \Gamma(\tau)} f_{\text{React}}^* d\tau d\sigma$$

for all control volumes $\Delta V$.

Interpreting $f_{\text{React}}^*$ as a surface production density, it means that for small $\epsilon$ the production on $\Omega_\varepsilon$ (here: reaction (1)) can be approximated by a reaction on $\Gamma$. Note that, under mild mathematical assumptions, this implies $\epsilon f_{\text{React}} \xrightarrow{\epsilon \to 0} f_{\text{React}}^*$. This fact yields information on the behaviour of the concentrations near $\Gamma$ of the species entering $f_{\text{React}}$, if one wants to approximate model $(P_{2\epsilon})$ by a model with $\Gamma(t)$-concentrated reactions. Such an attempt could be motivated by numerical advantages or others. The two cases mentioned above are $f_{\text{React}}^* = 0$ (no production on $\Gamma$) and $f_{\text{React}}^* \neq 0$.

Note that the solution output $\bar{X}_\varepsilon := (\bar{b}_\varepsilon, \bar{c}_\varepsilon, \bar{d}_\varepsilon, \bar{h}_\varepsilon, \bar{w}_\varepsilon, s_\varepsilon)$ also converges to a solution $\bar{X} := (\bar{b}, \bar{c}, \bar{d}, \bar{h}, \bar{w}, s)$ of the corresponding problem $(P_{2\epsilon=0})$ (with the volume rate $f_{\text{React}}$ replaced by the surface reaction rate $f_{\text{React}}^*$ (cf. Ref. [9]).

5 Simulation results for model $(P_{2\epsilon})$

5.1 Simulation strategy

In our simulations we will address the following questions:
1. How does a change of the structure of the reaction rate (exemplified by $p, q, r_1, r_2$ (cf. (13)) affects the output?

2. How does a change of the width $\epsilon$ of the main carbonation zone $\Omega_0$ affects the output?

3. How does a change of the weight of the reactions in $\Omega_0$ and $\Omega_0$, respectively, affects the output? Recall: A way of weighing is given by the weights $\delta_1$ and $\delta_2$ in (15). We will discuss several cases of $\delta = \frac{\delta_1}{\delta_2} \gg 1$ and $\delta \ll 1$.

4. Does the model reproduce some of the characteristics which are standard expectations for carbonation such as the $\sqrt{t}$-law for the penetration depth (for large $t$) or others?

5. Is the two-reaction-zones model ($P_{2\epsilon}$) capable of reproducing experimental results? More precisely: Let $r_1 = 0$ in (13), fix all data except $p$ and $q$ in (13) and $\alpha$ in (9). Are there reasonable $p, q, \alpha$ such that the simulation-output of model ($P_{2\epsilon}$) comes close to measurement data?

5.2 Outline of the simulation of model ($P_{2\epsilon}$)

Applying a fixing boundary technique [Landau (1950)] we transform the moving boundary problem ($P_{2\epsilon}$) into a problem with fixed boundaries. The price of this transformation is paid by obtaining space- and time-dependent coefficients in the transformed equations. By means of the weak formulation of the transformed problem together with a (piecewise linear) spline-based Galerkin ansatz, we obtain a system of $5n+1$ ode’s which we solve. Section 5.4 contains several numerical tests and their evaluation with respect to the chemical-physical mechanisms that usually appear in the carbonation of concrete-based materials.

5.3 Simulation procedure for ($P_{2\epsilon}$)

We make use of the change of variable

$$y = \begin{cases} \frac{x}{s(t) + \frac{\xi}{2}}, & \text{if } x \in [0, s(t) + \frac{\xi}{2}] \\ \frac{L-x}{L-s(t)-\frac{\xi}{2}}, & \text{if } x \in [s(t) - \frac{\xi}{2}, L] \end{cases}$$

(28)

to transform the domains $\Omega_0 \cup \Omega_0 \cap \Omega_0$ into the intervals $[l_\epsilon, 1]$ and $[0, 1]$, respectively, where $l_\epsilon = 1 - \frac{\epsilon}{s(t) + \frac{\xi}{2}}$ represents the image of the point $x = s(t) - \frac{\xi}{2}$ through the transformation (28).

After setting $c(y, t) = \tilde{c}(x, t)$, $d(y, t) = \tilde{d}(x, t)$ and $h(y, t) = \tilde{h}(x, t)$, and $b(y, t) = \tilde{b}(x, t)$, we homogenize the Dirichlet boundary conditions (24) prescribed in section 3.5. Using transformation (28), we obtain a system of equations on the fixed domain $0 < y < 1$. Therefore, the model ($P_{2\epsilon}$) (summarized in section 3.7) can be rewritten as:

$$c_{i,i} = \frac{D_{c_{i,i}}}{(s(t) + \frac{\xi}{2})^2} x c_{yy} + \frac{\hat{s}(t)}{s(t) + \frac{\xi}{2}} y c_{y} + f_{\text{Heny}} + \delta_{1} \chi_{[0, l_{\epsilon}]} f_{\text{Reac1}} + \delta_{2} \chi_{[l_{\epsilon}, 1]} f_{\text{Reac2}} + t > 0,$$  

(29)
\[
d_{t} = \frac{D_{d_{1}}}{(s(t) + \frac{\varepsilon}{T})^{2}} d_{yy} + \frac{\dot{s}(t)}{s(t) + \frac{\varepsilon}{T}} yd_{y} + f_{\text{H}_{1}\text{tr}} d_{t}, \quad t > 0, \tag{30}
\]

\[
h_{1,t} = \frac{D_{h_{1}}}{(s(t) + \frac{\varepsilon}{T})^{2}} h_{1,yy} + \frac{\dot{s}(t)}{s(t) + \frac{\varepsilon}{T}} yh_{1,y} + f_{h_{1}\text{tr}} d_{t} + \delta_{1} \chi_{(0,1)} f_{h_{1}\text{reac}} + \delta_{1} \chi_{(1,1)} f_{h_{1}\text{reac}}, \quad t > 0, \tag{31}
\]

\[
h_{2,t} = \frac{D_{h_{2}}}{(s(t) + \frac{\varepsilon}{T} - L)^{2}} h_{2,yy} + \frac{\dot{s}(t)}{s(t) + \frac{\varepsilon}{T} - L} yh_{2,y} + f_{h_{2}\text{tr}} d_{t}, \quad t > 0, \tag{32}
\]

\[
b_{t} = \frac{\dot{s}(t)}{s(t) + \frac{\varepsilon}{T}} yb_{y} + \delta_{1} \chi_{(0,1)} f_{b_{\text{reac}}}, \quad t > 0. \tag{33}
\]

Letting \(\bar{c}(y,0) = \lambda_{c}, \bar{d}(y,0) = \lambda_{d}, \bar{h}_{1}(y,0) = \lambda_{h_{1}}, \bar{h}_{2}(y,0) = \lambda_{h_{2}}\) and \(\bar{b}(y,0) = \lambda_{b}\), the initial conditions in 3.6 transform into:

\[
c(y,0) = 0, \quad d(y,0) = 0, \quad h_{1}(y,0) = 0, \quad h_{2}(y,0) = 0 \quad \text{and} \quad b(y,0) = 0. \tag{34}
\]

The boundary conditions associated to the model (3.2) (cf. sections 3.4, 3.5 and 3.6) become

\[
c(0,t) = 0, \quad -\frac{D_{c}}{s(t) + \frac{\varepsilon}{T}} c_{y}(1,t) = \dot{s}(t) (c(1,t) + \lambda_{c}), \quad t > 0, \tag{35}
\]

\[
d(0,t) = 0, \quad -\frac{D_{d}}{s(t) + \frac{\varepsilon}{T}} d_{y}(1,t) = \dot{s}(t) (d(1,t) + \lambda_{d}), \quad t > 0, \tag{36}
\]

\[
h_{1}(0,t) = 0, \quad \frac{D_{h_{1}}}{s(t) + \frac{\varepsilon}{T}} h_{1,y}(1,t) = \frac{D_{h_{2}}}{s(t) + \frac{\varepsilon}{T} - L} h_{2,y}(1,t) \quad \text{and} \quad h_{2}(0,t) = 0, \quad t > 0. \tag{37}
\]

Interface condition (24) turns into

\[
\dot{s}(t) = \frac{1}{\varepsilon} \frac{\int_{\Omega(t)} \tilde{n}_{k}(x,t) dx}{\int_{\Omega(t)} (h_{1}(x,t) + \lambda_{h_{1}}) dx}, \tag{38}
\]

where \(t > 0, \ 0 < y < 1\) and

\[
s(0) = s_{0}. \tag{39}
\]

To solve the system (29)-(39) numerically, we use its weak formulation and a (piecewise) linear-based Galerkin scheme. Let \(\{\psi_{j}^{n}\}_{j=0}^{n}\) denote the standard piecewise linear splines on the interval [0, 1] defined with respect to the uniform mesh \(\left\{\frac{i}{n}, \frac{i+1}{n}, \ldots, 1\right\}\). That is, for a fixed \(n \in \mathbb{N} \setminus \{0\}\) and \(j = 0, 1, 2, \ldots, n\), we introduce

\[
\psi_{j}^{n}(y) = \begin{cases} 
1 - |ny - j|, & \text{if } y \in \left[\frac{i}{n}, \frac{i+1}{n}\right] \cap [0, 1] \\
0, & \text{elsewhere on } [0, 1].
\end{cases}
\]
To obtain the Galerkin system, we set
\[c^n(y, t) = \sum_{j=1}^{n} C_j^n(t) \psi_j^n(y), \quad d^n(y, t) = \sum_{j=1}^{n} D_j^n(t) \psi_j^n(y),\]
\[h_k^n(y, t) = \sum_{j=1}^{n} H_{kj}^n(t) \psi_j^n(y), \quad b^n(y, t) = \sum_{j=1}^{n} B_j^n(t) \psi_j^n(y),\]
with \(k = 1, 2, t \geq 0, 0 \leq y \leq 1\) where
\[C^n(t) = [C_1^n(t), C_2^n(t), \ldots, C_n^n(t)]^T \in \mathbb{R}^n, \quad D^n(t) = [D_1^n(t), D_2^n(t), \ldots, D_n^n(t)]^T \in \mathbb{R}^n,
\]
\[H_k^n(t) = [H_{k1}^n(t), H_{k2}^n(t), \ldots, H_{kn}^n(t)]^T \in \mathbb{R}^n, \quad B^n(t) = [B_1^n(t), B_2^n(t), \ldots, B_n^n(t)]^T \in \mathbb{R}^n.\]
The superscript \(T\) stands for transposition.

The Galerkin equations are given by
\[M^n \dot{C}^n(t) = -\frac{D_1}{(s^n(t)+\frac{1}{2})} K^n C^n(t) + \frac{\dot{\psi}_1^n(t)}{s^n(t)+\frac{1}{2}} L^n C^n(t) - \frac{\dot{\psi}_1^n(t)}{s^n(t)+\frac{1}{2}} (C^n(t) + \lambda_{\varepsilon}) e^n + P_H (Q_H M^n D^n(t) - M^n C^n(t)) + P_H (Q_H \lambda_{\delta} - \lambda_0^\varepsilon) + \mu_1 f_{\varepsilon pure}(t) + \delta_c f_{\varepsilon pure}(t), t > 0,\]
\[M^n \dot{D}^n(t) = -\frac{D_1}{(s^n(t)+\frac{1}{2})} K^n D^n(t) + \frac{\dot{\psi}_1^n(t)}{s^n(t)+\frac{1}{2}} L^n D^n(t) - \frac{\dot{\psi}_1^n(t)}{s^n(t)+\frac{1}{2}} (D^n(t) + \lambda_{\delta}) e^n - P_H (Q_H M^n D^n(t) - M^n C^n(t)) - P_H (Q_H \lambda_{\delta} - \lambda_0^\varepsilon), t > 0,\]
\[M^n \dot{H}_1^n(t) = -\frac{D_1}{(s^n(t)+\frac{1}{2})} K^n H_1^n(t) + \frac{\dot{\psi}_1^n(t)}{s^n(t)+\frac{1}{2}} L^n H_1^n(t) - \frac{D_1}{(s^n(t)+\frac{1}{2})} (H_1^n(t) + \lambda_{\varepsilon}) e^n + \mu_1 f_{\varepsilon pure}(t) + \delta_c f_{\varepsilon pure}(t), t > 0,\]
\[M^n \dot{H}_2^n(t) = -\frac{D_2}{(s^n(t)+\frac{1}{2})} K^n H_2^n(t) + \frac{\dot{\psi}_1^n(t)}{s^n(t)+\frac{1}{2}} L^n H_2^n(t) - \frac{D_2}{(s^n(t)+\frac{1}{2})} (H_2^n(t) + \lambda_{\varepsilon}) e^n + \mu_1 f_{\varepsilon pure}(t) + \delta_c f_{\varepsilon pure}(t), t > 0,\]
\[M^n \dot{B}^n(t) = \frac{\dot{\psi}_1^n(t)}{s^n(t)+\frac{1}{2}} L^n B^n(t) + \mu_1 f_{\varepsilon pure}(t) + \delta_c f_{\varepsilon pure}(t), t > 0,\]

where \(f_{\varepsilon pure}(t) = f_{\varepsilon pure}(t) = -f_{\varepsilon pure}(t)\) and
\[f_{\varepsilon pure}(t) = \bar{\alpha} \frac{\mu_1}{\lambda_{\varepsilon max} \lambda_{\varepsilon max}} \int_0^1 (\dot{\psi}_1^n(t) - (\dot{b}(y, t) + \lambda_{\varepsilon}) r_1) (c^n(y, t) + \lambda_{\varepsilon}) (h_1^n(y, t) + \lambda_{\varepsilon}) r_2 \psi^n(y) dy,\]
with \(r_1 > 0, r_2 \in [0, 2]\) and
\[\bar{\alpha} := \alpha A e^{-\beta_\varepsilon \frac{\psi_1^n}{\psi_{\varepsilon pure}}} f_1(RH)\]

is a material parameter (cf. section 3.2.1).

Also, we set \(f_{\varepsilon pure}(t) = f_{\varepsilon pure}(t) = -f_{\varepsilon pure}(t)\), where
\[f_{\varepsilon pure}(t) = k \int_0^1 (c^n(y, t) + \lambda_{\varepsilon}) \psi^n(y) dy, t > 0.\]
The interface relation (38) becomes:

\[
\delta^{\alpha}(t) = \kappa e \int_{I_{10}}^{1} \left( e^{\alpha}(y, t) + \lambda^{\alpha}_{1}(y, t) \right)^{q} dy \\
\int_{I_{10}}^{1} \left( h^{\alpha}_{1}(y, t) + \lambda^{\alpha}_{1} \right) dy, \tag{43}
\]

where \( t > 0, p, q \in [0, 2] \), and

\[
s^{\alpha}(0) = s_{0} \geq d_{10} + \frac{\varepsilon}{2} > 0. \tag{44}
\]

d_{10} denotes the width\(^3\) of the initial layer \( \Omega_{10} \). See the reference picture Fig. 6.

The initial conditions are

\[
C^{\alpha}(0) = 0, D^{\alpha}(0) = 0, H^{\alpha}_{1}(0) = 0, H^{\alpha}_{2}(0) = 0 \text{ and } B^{\alpha}(0) = 0. \tag{45}
\]

**Remark 1** The matrices \( M^{\alpha}, K^{\alpha} \in \mathbb{R}^{(n+1) \times (n+1)}, L^{\alpha}_{i} \in \mathbb{R}^{n \times n} \) and \( e^{1}, e^{n}, \lambda^{\alpha}_{e}, \lambda^{\alpha}_{d} \in \mathbb{R}^{n} \), \( n \in \mathbb{N}, n > 1 \) are given by

\[
[M^{\alpha}]_{ij} = \int_{0}^{1} \psi_{i}^{\alpha}(y) \psi_{j}^{\alpha}(y) dy, \quad [K^{\alpha}]_{ij} = \int_{0}^{1} \psi_{i}^{\alpha r}(y) \psi_{j}^{\alpha r}(y) dy, \quad i, j = 1, 2, \ldots, n + 1,
\]

\[
[L^{\alpha}_{i}]_{ij} = \int_{0}^{1} y \psi_{i}^{\alpha}(y) \psi_{j}^{\alpha r}(y) dy, \quad i, j = 1, 2, \ldots, n
\]

and

\[
e^{1} = [1, 0, \ldots, 0, 0]^{T}, \quad e^{n} = [0, 0, \ldots, 0, 1]^{T},
\]

\[
\lambda^{\alpha}_{e} = \lambda^{\alpha}_{e} \left[ \frac{1}{n}, \frac{1}{n}, \ldots, \frac{1}{n}, \frac{1}{2n} \right]^{T}, \quad \lambda^{\alpha}_{d} = \lambda^{\alpha}_{d} \left[ \frac{1}{n}, \frac{1}{n}, \ldots, \frac{1}{n}, \frac{1}{2n} \right]^{T}.
\]

The stiffness matrix \( M^{\alpha} \) is sparse, symmetric, positive definite and diagonally dominant. We note that \( cond(M^{\alpha}) \approx 3.9938 \) for \( n = 32 \) and \( cond(M^{\alpha}) \approx 3.9993 \) for \( n = 100 \). We used MATLAB routine \texttt{ode15s} to integrate the system of ode’s (41), (43) with the initial values (44), (45).

\(^2\)The factor \( \kappa \) is given in Ref. [29]) accounts for the effect of the variation of exponents \( p, q \) on the carbonation rate constant \( k \). Since we are not aware of experimental data for such effect, a (possible) threshold value for this parameter is found numerically by variation of the value of an auxiliary parameter \( \alpha_{1} > 0 \). Once better measurements of carbonation rate constant are available, the parameter \( \alpha_{1} \) can be identified.

\(^3\)In the simulations we chose \( s_{0} = d_{10} + \frac{\varepsilon}{2} \) with \( d_{10} = 10^{-1} \).
5.4 Simulation curves

Figure 7: (a) Profiles of $CO_2(g)$ in $\Omega_1(t) \cup \Omega_\varepsilon(t)$. One observes that the exposure concentration of $CO_2(g)$ was split via Henry’s law in the initial layer. $CO_2$ is strongly consumed in $\Omega_\varepsilon(t)$. (b) Profiles of $Ca(OH)_2$ in $\Omega_2(t)$. $(D_{d_1}, k, \alpha_1, p, q, r_1, r_2, \varepsilon) = (3.5, 5 \times 10^3, 10^2, 2, 0.5, 1, 1, \frac{1}{100})$ and $\frac{\delta_0}{\delta_1} \ll 1$.

Figure 8: Carbonation penetration (in cm) after 9 weeks of accelerated testing. The graphs were obtained varying $\delta_1 = \frac{\delta_0}{2}, \frac{\delta_0}{4}, \frac{\delta_0}{6}, \frac{\delta_0}{8}, \frac{\delta_0}{10}$, where $\delta_0 = 10^{-5}\delta_c$. $(k, D_{d_1}, p, q, \alpha_1, r_1, r_2, \varepsilon) = (5 \times 10^3, 3.5, 2, 0.5, 10^2, 1, 1, \frac{1}{100})$ in the case $\frac{\delta_0}{\delta_1} \gg 1$. 


Figure 9: \( \text{CaCO}_3 \) concentration in \( \Omega_1(t) \cup \Omega_2(t) \) after 9 weeks of accelerated testing. One observes that after each time interval (7 days) some calcium carbonate is produced. This production stops when all available \( \text{Ca(OH)}_2 \) in \( \Omega_1(t) \cup \Omega_2(t) \) is consumed. Here \( (k, D_{d1}, p, q, \alpha, r_1, r_2, \epsilon) = (5 \times 10^3, 3.5, 2, 0.5, 10^2, 1, 1, \frac{1}{100}) \) and \( \frac{\delta}{\delta_1} \gg 1 \).

Figure 10: \( \text{CaCO}_3 \) concentration in \( \Omega_1(t) \cup \Omega_2(t) \) after 9 weeks of accelerated testing. Here \( (k, D_{d1}, p, q, \alpha, r_1, r_2, \epsilon) = (5 \times 10^3, 3.5, 2, 0.9, 10^2, 1, 1, \frac{1}{100}) \) in the case \( \frac{\delta}{\delta_1} \gg 1 \). One can remark smoothing effects produced on the concentration profiles by the introduction of an artificial diffusive flux of \( \text{CaCO}_3 \) (aq) with \( D_{b1} = 10^{-4} \).
Figure 11: CaCO$_3$ concentration in $\Omega_1(t) \cup \Omega_2(t)$ after 9 weeks of accelerated testing. Here we used the same parameters as in Fig 9: $(k, D_{d_1}, p, q, \alpha_1, r_1, r_2, \epsilon) = (5 \times 10^3, 3.5, 2, 0.5, 10^2, 1, 1, \frac{1}{10})$ but with a bigger ratio $\frac{\delta_1}{\delta_2} \gg 1$ (more precisely, with a smaller $\delta_1$).

Figure 12: CaCO$_3$ concentration in $\Omega_1(t) \cup \Omega_2(t)$ after 9 weeks of accelerated testing. Here $(k, D_{d_1}, p, q, \alpha_1, r_1, r_2, \epsilon, d_{d_0}) = (5 \times 10^3, 3.5, 2, 0.9, 10^2, 1, 1, \frac{1}{10}, 0.1)$ in the case $\frac{\delta_1}{\delta_2} \gg 1$. One can remark smoothing effects produced on the concentration profiles by the introduction of an artificial diffusive flux of CaCO$_3$(aq) with $D_{b_1} = 10^{-4}$. 
Figure 13: Carbonation penetration (in cm) after 9 weeks of accelerated testing. The graphs were obtained varying the exponents \((p, q) = (2, 0.5), (2, 0.7), (2, 0.9)\). Here \((k, D_{d_1}, \alpha_1, r_1, r_2, \epsilon) = (5 \times 10^3, 3.5, 10^2, 1, 1, \frac{1}{100})\) and \(\frac{\delta}{d_1} \gg 1\).

Figure 14: Carbonation penetration (in cm) after 9 weeks of accelerated testing. The graphs were obtained varying the diffusivity \(D_{d_1} = \frac{14}{8}, \frac{21}{8}, \frac{28}{8}\). Here \((k, p, q, \alpha_1, r_1, r_2, \epsilon) = (5 \times 10^3, 1.8, 0.8, 10^2, 1, 1, \frac{1}{100})\) and \(\frac{\delta}{d_1} \gg 1\).
Figure 15: Carbonation penetration (in cm) after 9 weeks of accelerated testing. The graphs were obtained varying $\epsilon = \frac{1}{25}, \frac{1}{50}, \frac{1}{100}, \frac{1}{150}, \frac{1}{300}$. Here $(k, D_{d_1}, p, q, \alpha_1, r_1, r_2) = (5 \times 10^3, 3.5, 2, 0.8, 10^2, 1, 1)$ and $\frac{2}{\epsilon_1} \gg 1$. One observes that the bigger the $\epsilon$ is, the faster the speed becomes. $\epsilon = \frac{1}{300}$ leads to numerical instability.

Figure 16: Carbonation penetration (in cm) after 9 weeks of accelerated testing. The graphs were obtained varying $r_2 = 0, 0.5, 0.75, 1, 1.5$. The blow up appears when $r_2 = 1.5$. $(k, D_{d_1}, p, q, \alpha_1, r_1, \epsilon) = (5 \times 10^3, 3.5, 1.8, 1.5, 10^2, 1, \frac{1}{100})$ and $\frac{2}{\epsilon_1} \gg 1$. 
Figure 17: $CaCO_3$ concentration in $\Omega_1(t) \cup \Omega_2(t)$ after 9 weeks of accelerated testing. Here $(k, D_{d_1}, p, q, \alpha_1, r_1, r_2, \epsilon) = (5 \times 10^3, 3.5, 2, 0, 10^2, 1, 1, \frac{1}{8})$ and $\frac{\epsilon}{d_1} \ll 1$. Here $CaCO_3(aq)$ is allowed to diffusive very slowly ($D_{b_1} = 10^{-4}$).

Figure 18: $CaCO_3$ concentration in $\Omega_1(t) \cup \Omega_2(t)$ after 9 weeks of accelerated testing. Here $(k, D_{d_1}, p, q, \alpha_1, r_1, r_2, \epsilon) = (5 \times 10^3, 3.5, 2, 0, 10^2, 1, 1, \frac{1}{8})$ and $\frac{\epsilon}{d_1} \ll 1$. 
5.5 Summary and conclusions

5.5.1 Interpretations of the simulation results

1. The model shows several general characteristics one might expect:

   (a) Penetration-depth curves exhibit a near $t^{1/2}$-behavior for large $t$ (cf. Fig. 8, Fig. 13 - Fig. 16). Note: In the context of accelerated carbonation 6 to 9 weeks is large!

   (b) $CO_2$-curves correspond to results known from other models.

   (c) A large $Ca(OH)_2$-concentration gradient in $\Omega_2$ near $\Omega_c$ can be observed (cf. Fig. 7 (b)).

2. The model shows effects based on the two-zone- and moving-boundary setting:

   (a) The (example of) $Ca(OH)_2$-concentration profiles over several weeks (cf. Fig. 7 (b)) shows the effect of an accentuated ($\delta \gg 1$) influence of the reaction in $\Omega_2$. Although the model provides for an (indefinite) re-production of $Ca(OH)_2$ by dissolution, the consumption due to reaction (1) is much stronger than in $\Omega_1$. This effect can be controlled quantitatively by the weighing parameter $\delta$.

   (b) One of the motivation for the introducing a two-reaction-zones model has been the experimental observation, that after a fast increase of the carbonation degree near the front a slow (with respect to time) gradual increase occurs in $\Omega_1$. This behavior of the model output is shown in Fig. 9 - Fig. 12, Fig. 17 and Fig. 18. A conjecture is, that the speed of this increase can be controlled by an appropriate (but not yet known) combination of $\delta_1$ and $\delta_c$ and the size of the reaction constant for the reaction on $\Omega_1$.

   (c) The interplay between the two reaction zones is reflected by the (slight) increase of the $CaCO_3$ curve in the main-reaction zone $\Omega_c$ if $\delta \gg 1$ (cf. Fig. 9 - Fig. 12). For $\delta \ll 1$ one obtains a slight decrease in $\Omega_c$ (cf. Fig. 17 and Fig. 18).

3. A classification for the behavior of the penetration-depth curves with respect to $p, q, r_1$ and $r_2$ (cf. introduction of the reaction rates in (9), (12)) is open. In particular, one needs more information on which of these parameters are the more relevant ones. For a comprehensive model possibly modeling the carbonation of $CSH$-phases, too, an approach using too many parameters will be useless due to the lack of data to fit these parameters.

   By a mere variation of $p, q, r_1$ and $r_2$, respectively, one obtains a large variety of carbonation-penetration depth curves and concentration profiles (cf. Fig. 8, Fig. 13 - Fig. 16).

5.5.2 Conclusions

The two-reaction-zone model presented in this note shows several of the major characteristics expected from a carbonation-prediction model (shape of the curves, magnitude of the curves for real-material data). The model output is stable with respect to most of the parameters. The large variety of penetration-curves under a change of some of the
reaction-rates parameters indicates the necessity of a reduction of the number of reaction-rate parameters. Moreover, the introduction of two reaction zones (with possibly-but not necessarily) different reaction rates allows a more detailed analysis of the carbonation behavior near and far from the reaction front and how this behavior can model-wise be controlled.

6 Appendix

<table>
<thead>
<tr>
<th>Quantity</th>
<th>Definition</th>
<th>Dimensions</th>
<th>Value(range)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$D_{w1}, D_{w2}$</td>
<td>Effective moisture diffusivities, Refs. [46, 26]</td>
<td>cm$^2$ day$^{-1}$</td>
<td>[0.9, 90]</td>
</tr>
<tr>
<td>$D_h$</td>
<td>Effective $Ca(OH)_2$(aq) diffusivity, Ref. [35]</td>
<td>cm$^2$ day$^{-1}$</td>
<td>0.864</td>
</tr>
<tr>
<td>$D_{c1}$</td>
<td>Effective $CO_2$(aq) diffusivity, Ref. [11]</td>
<td>cm$^2$ day$^{-1}$</td>
<td>[0.62, 6.2]</td>
</tr>
<tr>
<td>$\lambda_{m1}$</td>
<td>Initial values of moisture $i=1,2$, Table 5</td>
<td>g cm$^{-3}$</td>
<td>[0.061, 0.123]</td>
</tr>
<tr>
<td>$\lambda_l$</td>
<td>Initial concentration of $CO_2$(g), Ref. [45]</td>
<td>g cm$^{-3}$</td>
<td>$58.92 \times 10^{-4}$</td>
</tr>
<tr>
<td>$\lambda_i$</td>
<td>Initial value for $CaCO_3$</td>
<td>g cm$^{-3}$</td>
<td>0</td>
</tr>
<tr>
<td>$\lambda_r$</td>
<td>Initial value for $Ca(OH)_2$(aq), Ref. [26]</td>
<td>g cm$^{-3}$</td>
<td>$77.5 \times 10^{-3}$</td>
</tr>
<tr>
<td>$s_0$</td>
<td>Initial position cf. Fig. 6</td>
<td>cm</td>
<td>10$^{-5}$</td>
</tr>
<tr>
<td>$2L$</td>
<td>Length of the observed slab, Ref. [45]</td>
<td>cm</td>
<td>10</td>
</tr>
<tr>
<td>$\phi_1$</td>
<td>Porosity of non-carbonated concrete, Ref. [26]</td>
<td>-</td>
<td>0.15</td>
</tr>
<tr>
<td>$\phi_2$</td>
<td>Porosity of carbonated concrete, Ref. [26]</td>
<td>-</td>
<td>0.13</td>
</tr>
<tr>
<td>$Q_H$</td>
<td>Exchange term in Henry's law, Refs. [35, 6]</td>
<td>-</td>
<td>0.8227</td>
</tr>
<tr>
<td>$P_H$</td>
<td>Mass transfer coefficient of $CO_2$(g) in pore water, Ref. [6]</td>
<td>day$^{-1}$</td>
<td>35760</td>
</tr>
<tr>
<td>$k_T$</td>
<td>Mass transfer constant of $CO_2$(g), air to water, Ref. [6]</td>
<td>cm day$^{-1}$</td>
<td>7</td>
</tr>
<tr>
<td>$P_{hk\text{diss}}$</td>
<td>Factor in the dissolution law, $k=1,2$</td>
<td>day$^{-1}$</td>
<td>$\frac{1}{196}$</td>
</tr>
<tr>
<td>$\Delta S$</td>
<td>Surface area to volume ratio, for air, Ref. [6]</td>
<td>cm$^{-1}$</td>
<td>$10^4$</td>
</tr>
<tr>
<td>$\Delta S_{\text{air}}$</td>
<td>Surface area to volume ratio, for water, Ref. [6]</td>
<td>cm$^{-1}$</td>
<td>$10^4$</td>
</tr>
</tbody>
</table>

Table 1: Numerical data for several parameters and input variables.

\*The threshold values introduced in this table are taken from the literature. We use numerical ranges and not precise values since we are not aware of measurements for these parameters within the ace in Ref. [45].

<table>
<thead>
<tr>
<th>Quantity</th>
<th>Definition</th>
<th>Dimensions</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$R$</td>
<td>Gas constant, Ref. [13]</td>
<td>mol$^{-1}$ K$^{-1}$ atm</td>
<td>$8206 \times 10^{-5}$</td>
</tr>
<tr>
<td>$H$</td>
<td>Henry's law constant for $CO_2$(g), Ref. [36]</td>
<td>mol m$^{-3}$ atm$^{-1}$</td>
<td>34.2</td>
</tr>
<tr>
<td>$M_{H_2O}$</td>
<td>Molecular weight of water, Ref. [13]</td>
<td>g mol$^{-1}$</td>
<td>18</td>
</tr>
<tr>
<td>$M_{CO_2}$</td>
<td>Molecular weight of $CO_2$, Ref. [13]</td>
<td>g mol$^{-1}$</td>
<td>44</td>
</tr>
<tr>
<td>$M_{CaCO_3}$</td>
<td>Molecular weight of aragonite/calcite, Ref. [13]</td>
<td>g mol$^{-1}$</td>
<td>100.087</td>
</tr>
<tr>
<td>$M_{Ca(OH)_2}$</td>
<td>Molecular weight of $Ca(OH)_2$, Ref. [13]</td>
<td>g mol$^{-1}$</td>
<td>74</td>
</tr>
<tr>
<td>$\rho_{Ca(OH)_2}$</td>
<td>Density of $Ca(OH)_2$, Ref. [13]</td>
<td>g cm$^{-3}$</td>
<td>2.24</td>
</tr>
<tr>
<td>$\rho_{CaCO_3}$</td>
<td>Density of $CaCO_3$ (calcite), Ref. [13]</td>
<td>g cm$^{-3}$</td>
<td>2.71</td>
</tr>
</tbody>
</table>

Table 2: Useful physical and material constants.
<table>
<thead>
<tr>
<th>Quantity</th>
<th>Definition</th>
<th>Dimensions</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$w/c$</td>
<td>Water:cement ratio</td>
<td>-</td>
<td>0.60</td>
</tr>
<tr>
<td>$a/c$</td>
<td>Aggregate:cement ratio</td>
<td>-</td>
<td>5.1429</td>
</tr>
<tr>
<td>$\rho_c$</td>
<td>Cement density, Refs. [45, 20]</td>
<td>g cm$^{-3}$</td>
<td>3.15</td>
</tr>
<tr>
<td>$\rho_a$</td>
<td>Aggregate density, Ref. [45]</td>
<td>g cm$^{-3}$</td>
<td>2.7</td>
</tr>
<tr>
<td>$\rho_{H_2O}$</td>
<td>Water density, Refs. [45, 20]</td>
<td>g cm$^{-3}$</td>
<td>1</td>
</tr>
<tr>
<td>$\rho_{cp}$</td>
<td>Cement paste density (CEM I 0.60), Ref. [45]</td>
<td>g cm$^{-3}$</td>
<td>1.7439</td>
</tr>
<tr>
<td>$\rho_{CaO}$</td>
<td>Calcium oxide density, Ref. [13]</td>
<td>g cm$^{-3}$</td>
<td>3.34</td>
</tr>
</tbody>
</table>

Table 3: Material characteristics of the concrete sample.

<table>
<thead>
<tr>
<th>Portland cement (CEM I)</th>
<th>RH (%)</th>
<th>$\bar{w}$ (kg/m$^3$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$SiO_2$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$Al_2O_3$</td>
<td>20</td>
<td></td>
</tr>
<tr>
<td>$Fe_2O_3$</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>$CaO$</td>
<td>63</td>
<td></td>
</tr>
<tr>
<td>$MgO$</td>
<td>1.5</td>
<td></td>
</tr>
<tr>
<td>$SO_3$</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>$Na_2O$ and $K_2O$</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Others</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Loss on ignition</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Insoluble residue</td>
<td>0.5</td>
<td></td>
</tr>
</tbody>
</table>

Table 4: Composition (mass fraction %), cf. Ref. [27].

Table 5: Material data for concrete showing the relation between relative humidity RH and moisture content $\bar{w}$, cf. Ref. [1], III.4. See also Fig. 4 in Ref. [22].

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